

This is a repository copy of *Connecting Earth Observation to High-Throughput Biodiversity Data*.

White Rose Research Online URL for this paper:

<https://eprints.whiterose.ac.uk/118918/>

Version: Accepted Version

Article:

White, Piran Crawford Limond orcid.org/0000-0002-7496-5775 and Yu, Douglas (2017) Connecting Earth Observation to High-Throughput Biodiversity Data. *Nature Ecology and Evolution*. pp. 1-9.

<https://doi.org/10.1038/s41559-017-0176>

Reuse

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.

1 Connecting Earth Observation to High-Throughput

2 Biodiversity Data

3 Bush, Alex^{1,2,3}; Sollmann, Rahel⁴; Wilting, Andreas⁵; Bohmann, Kristine^{6,7}; Cole, Beth⁸;
4 Balzter, Heiko^{8,9}; Martius, Christopher¹⁰; Zlinszky, András¹¹; Calvignac-Spencer,
5 Sébastien¹²; Cobbold, Christina A.¹³; Dawson, Terence P.¹⁴; Emerson, Brent C.^{15,7};
6 Ferrier, Simon³; Gilbert, M. Thomas P.^{6,16}; Herold, Martin¹⁷; Jones, Laurence¹⁸;
7 Leendertz, Fabian H.¹²; Matthews, Louise¹³; Millington, James D.A.¹⁴; Olson, John R.¹⁹;
8 Ovaskainen, Otso^{20,21}; Raffaelli, Dave²²; Reeve, Richard¹³; Rödel, Mark-Oliver²³;
9 Rodgers, Torrey W.²⁴; Snape, Stewart²⁵; Visseren-Hamakers, Ingrid²⁶; Vogler, Alfried
10 P.^{27,28}; White, Piran C.L.²²; Wooster, Martin J.¹⁴; Yu, Douglas W.^{1,7, †}

11

12 1. State Key Laboratory of Genetic Resources and Evolution, **Kunming Institute of Zoology**, Chinese

13 Academy of Sciences, Kunming, Yunnan 650203 China

14 2. Environment Canada @ Canadian Rivers Institute, Department of Biology, **University of New**

15 **Brunswick**, PO Box 4400, Fredericton, NB, E3B 5A3, Canada

16 3. **CSIRO Land and Water**, Canberra, ACT 2601, Australia

17 4. Department of Wildlife, Fish, & Conservation Biology, 1088 Academic Surge, One Shields Ave,

18 **University of California Davis**, Davis, CA 95616 USA

19 5. **Leibniz Institute for Zoo and Wildlife Research**, Alfred-Kowalke-Str. 17, 10315 Berlin, Germany

- 20 6. EvoGenomics, Natural History Museum of Denmark, **University of Copenhagen**, 1350 Copenhagen
- 21 K Denmark
- 22 7. School of Biological Sciences, **University of East Anglia**, Norwich Research Park, Norwich, Norfolk
- 23 NR47TJ UK
- 24 8. Centre for Landscape and Climate Research and Leicester Institute for Space and Earth Observation
- 25 (LISEO), **University of Leicester**, University Road, Leicester LE17RH UK
- 26 9. NERC National Centre for Earth Observation (NCEO) at **University of Leicester**, University Road,
- 27 Leicester LE17RH, UK
- 28 10. **Center for International Forestry Research (CIFOR)**, P.O. Box 0113 BOCBD, Bogor 16000,
- 29 Indonesia
- 30 11. Balaton Limnological Institute, **Centre for Ecological Research**, Hungarian Academy of Sciences,
- 31 Tihany 8237 Hungary
- 32 12. **Robert Koch Institut**, Berlin 13353 Germany
- 33 13. Boyd Orr Centre for Population and Ecosystem Health, **University of Glasgow**, Glasgow G128QQ UK
- 34 14. Department of Geography, **King's College London**, Strand Campus, London WC2R2LS UK
- 35 15. **IPNA-CSIC**, La Laguna, Tenerife, Canary Islands 38206 Spain
- 36 16. NTNU University Museum, **Norwegian University of Science and Technology**, Trondheim 7491
- 37 Norway
- 38 17. Laboratory of Geo-Information Science and Remote Sensing, **Wageningen University & Research**,
- 39 Wageningen, the Netherlands
- 40 18. Centre for Ecology and Hydrology, **Environment Centre Wales**, Deiniol Road, Bangor LL572UW UK
- 41 19. Division of Hydrologic Sciences, **Desert Research Institute**, Las Vegas, NV, USA
- 42 20. Department of Biosciences, **University of Helsinki**, Helsinki FI-00014 Finland

- 43 21. Centre for Biodiversity Dynamics, Department of Biology, **Norwegian University of Science and**
44 **Technology**, Trondheim 7491 Norway
- 45 22. Environment Department, **University of York**, York YO105NG UK
- 46 23. Museum für Naturkunde - **Leibniz Institute for Evolution and Biodiversity Science**, Berlin 10115
47 Germany
- 48 24. Department of Wildland Resources, **Utah State University**, Logan, UT 84322 USA
- 49 25. **Forestry Commission**, Edinburgh, UK
- 50 26. Department of Environmental Science and Policy, **George Mason University**, Fairfax, VA 22030 USA
- 51 27. Department of Life Sciences, **Natural History Museum**, London SW75BD UK
- 52 28. Department of Life Sciences, Silwood Park Campus, **Imperial College London**, Ascot SL57PY UK
- 53 ‡ Corresponding author: dougwyu@gmail.com

54 **Preface**

55 There is much interest in using Earth Observation (EO) technology to track biodiversity,
56 ecosystem functions, and ecosystem services, understandable given the fast pace of
57 biodiversity loss. However, because most biodiversity is invisible to EO, EO-based
58 indicators could be misleading, which can reduce the effectiveness of nature
59 conservation and even unintentionally decrease conservation effort. We describe an
60 approach that combines automated recording devices, high-throughput DNA
61 sequencing, and modern ecological modelling to extract much more of the information

62 available in EO data. This approach is achievable now, offering efficient and near-real-
63 time monitoring of management impacts on biodiversity and its functions and services.

64 **Meeting the Aichi Biodiversity Targets**

65 From Google Earth to airborne sensors, the Copernicus Sentinels, and cube satellites,
66 Earth Observation is undergoing a rapid expansion in capacity, accessibility, resolution,
67 and signal-to-noise ratio, resulting in a recognised shift in our capability for using
68 remote-sensing technologies to monitor biophysical processes on land and water¹⁻³.
69 These advances are motivating calls to use Earth Observation products to manage our
70 natural environment and to track progress toward global and national policy targets on
71 biodiversity and ecosystem services⁴⁻⁶. Foremost among these policies are the Strategic
72 Plan for Biodiversity and the Aichi Biodiversity Targets, which were adopted in 2010 by
73 the Parties to the Convention on Biological Diversity (CBD) to "take effective and urgent
74 action to halt the loss of biodiversity in order to ensure that by 2020 ecosystems are
75 resilient and continue to provide essential services..."⁷. The United Nations Sustainable
76 Development Goals⁸ now include some of the Aichi Targets, and the 2015 Paris
77 Agreement has reiterated the commitments of the UN Framework Convention on
78 Climate Change to reducing emissions from deforestation and forest degradation

79 (REDD+) and to securing non-carbon benefits, which include biodiversity and ecosystem
80 services⁹.

81 However, we have struggled to track and report progress toward the Aichi Targets in a
82 standardised and comprehensive way¹⁰. Although almost two-thirds of the CBD Parties
83 have updated their National Biodiversity Strategies and Action Plans to reflect the 2010
84 revisions, many still do not contain measurable indicators on the state of biodiversity, let
85 alone ecosystem services. This lack of quantification conceals the impacts of policy and
86 management interventions on biodiversity and ecosystem functions and services¹¹. The
87 difficulty of designing indicators¹²⁻¹⁴ has prompted an international consortium of
88 biodiversity scientists called GEO BON (Group on Earth Observations' Biodiversity
89 Observation Network) to propose a framework of Essential Biodiversity Variables¹⁵, with
90 the aim of setting minimum standards of coverage to ensure informativeness and to
91 harmonise disparate local measures so that biodiversity and ecosystem data can be
92 compared over space and time. The Essential Biodiversity Variables thus measure the
93 'state of biodiversity' at multiple levels: genetic composition, species populations,
94 species traits, community composition, ecosystem structure, and ecosystem function¹⁵.

95 Although it was originally envisioned that most of the variables (genetic to community
96 composition) would be scaled up from "intensive *in-situ* measurements"¹⁵ taken on the

97 ground, such measurements are costly and difficult because they are traditionally
98 gathered by visual and aural detection of plants and animals in the wild (preceded by
99 months or years of observer practice) and by mass collection of organisms (followed by
100 months of identification from morphology), so that data collection is slowed by human-
101 caused bottlenecks in sampling and taxonomy¹⁶.

102 As a result, attention is now being focused on designing 'Satellite Remote Sensing-
103 Essential Biodiversity Variables' (SRS-EBVs) to enable cost-effective and global-scale
104 monitoring^{5,6,12}. The problem here is that only a few Earth Observation products can be
105 mapped directly to Essential Biodiversity Variables and then to Aichi Targets, because
106 these products primarily measure gross vegetation and landscape metrics, such as land
107 cover and phenology⁴. For example, Pettorelli et al.¹² found only two Earth Observation
108 products (net primary productivity and fire incidence) that could serve as Essential
109 Biodiversity Variables for the Sahara, despite this biome's suitability for remote sensing
110 due to its visible biodiversity hotspots, remoteness, and availability of long time series.
111 Many of the Aichi Targets require data with species-level resolution, either because some
112 species are direct policy targets (e.g. Target 9: "invasive species controlled or eradicated")
113 or because species compositional data define the metric (e.g. Target 11: "protected areas
114 are ecologically representative and conserved effectively").

Clearly, a radically new approach is required if progress towards the Aichi Targets is to be accelerated, one that is robust, widely affordable, and can record stocks and changes in biodiversity and ecosystem services consistently, continuously, and at high resolution over large geographic scales. Here, we present such an approach in a framework that exploits recent efficiency gains and analytical breakthroughs in sensors, computation, ecology, taxonomy, and genomics (**Figure 1, Box 1**).

Box 1. Inferring a Hidden Ecosystem Function from Space

Large-bodied Amazonian monkeys are responsible for a key ecosystem function: they are the primary dispersers of large seeds, which are associated with more carbon-dense tree species. Peres et al.¹⁷ have proposed that this function boosts forest carbon storage. The idea can be tested by using Earth Observation data and public records to map human settlements and transport corridors and predict where monkey populations have declined through hunting^{17,18}. We can then use on-the-ground sampling and airborne sensors to test whether forests that have had longer exposure to hunting lack monkey populations and have more low-carbon-density tree species dispersed by wind and birds. In short, by combining Earth-Observation-derived maps of human activity with empirical observations of the response of primate populations to that activity, it should be

possible to map and track an ecosystem function (large-seed dispersal) that is invisible to satellites but contributes to an important ecosystem service (climate regulation).

From Point Samples to Continuous Maps

Instead of trying to map Earth Observation (EO) products directly to biodiversity, as encapsulated by SRS-EBVs^{4-6,12}, we propose to extract more information from EO data by interpolating biodiversity point samples to build continuous landscape maps of species distributions (**Figure 1**)¹⁹. Because it is species that are mapped, it then becomes possible to layer on the vast biological knowledge that we have collectively built up over decades of research, including historical distributions, phylogenetic relationships, and knowledge of species traits and interactions to infer, map, and track the distributions of ecosystem functions and services (**Box 1**). This approach, which we call here **CEOBE** (Connecting Earth Observation to Biodiversity and Ecosystems), is possible because of (1) major advances in EO sensitivity and capacity, (2) more efficient techniques to collect biodiversity data on the ground, and (3) modern community-analysis models from statistical ecology. We now review each of these advances, with additional detail in Supplementary Information.

The New Era of Earth Observation

There are ten times as many satellites in operation now as there were in the 1970s, a result of increasing sensor longevity and a six-fold increase in launches²⁰. Spatial resolution has improved to less than 1 m in both optical and radar sensors. Data continuity is also being maintained, most directly by the launch of NASA's Landsat 8 in 2013, which extends and technically enhances the 40-year Landsat record of medium-resolution, multispectral surface observations²¹. Data continuity is a key factor in understanding changes in biodiversity, as threats to biodiversity impact at a range of scales and often across lengthy timespans²².

The long-term Landsat record is being enhanced by new satellite systems and multiple sensors in a global network, a 'virtual constellation' that may help overcome problems in terrestrial monitoring from single sensors². As part of the Copernicus program, the ESA Sentinel satellites are the latest addition to the global network. With six missions planned and the first three launched, the Sentinels have radar, optical sensors, radiometers, and spectrometers with different goals²³. Sentinel-1, the radar satellite, and Sentinel-2, the superspectral high-resolution mission, are of particular interest to biodiversity monitoring, with long-term continuity of measurements, global coverage, and quick revisit times^{24,25}.

170 There have also been developments in hyperspectral sensors with EnMAP, HypsIRI,
171 PRISMA, and FLEX imaging spectrometer missions planned¹. In addition, airborne data
172 collection using high-resolution 3D airborne laser scanning is complementing spectral
173 information with structure²⁶. Swarms of commercial cube satellites and the use of drones
174 to carry sensors are additional significant steps that complement these large-scale
175 programs (**Supplementary Note 1** "Earth Observation technology").

176 The increase in spatial resolution in the new sensors implies greater precision because
177 reference measurements taken within meter-scale plots on the ground can be matched
178 directly to meter-scale pixels²⁷. This in turn improves the ability of EO to recognise
179 spatial gradients and boundaries.

180 Two additional factors affect the utility of remote sensing data for understanding
181 biodiversity change (**Supplementary Note 2** "Biodiversity and ecosystem information in
182 EO data"): affordability and access²². There has been a cultural shift, with free open
183 access on the rise. The opening of the Landsat archive in 2008 was a monumental
184 development²⁸, with ESA's Copernicus program following suit. Data access also refers to
185 the ability of users to retrieve, manipulate, and extract value from EO data. Cloud
186 computing and toolboxes are making these processes manageable, even with large data
187 archives.

The availability of copious EO data that have been shown in multiple studies to correlate closely with on-the-ground measures of ecosystem structure, habitat condition, and even animal communities (**Supplementary Note 2**) might suggest that remote sensors can be used directly to define environmental indicators, but we must acknowledge that we are still in the early stages of understanding how biodiversity delivers ecosystem functions and services, and how they all respond to exogenous change. Directly observing functional diversity is a partial solution but only with visible biodiversity such as vegetation²⁶. Thus, the challenge is to find ways to exploit the high efficiency and information content of EO data while not falling prey to *reification fallacy* (**Box 2**), which can arise when convenient but incomplete indicators are made available^{29,30}. Our institutions and reporting systems then retain the option to add and respond to new knowledge.

Box 2. The Perils of Convenient Indicators

If we rely too directly on EO data, we run the risk of *reification fallacy*, in which a mere indicator of a policy target itself ends up the target. Reification fallacy can reduce or narrow conservation effort³¹ and can crowd out future discoveries³². For example, while remote sensing is an efficient and direct way to measure forest *cover* (Aichi Target 5:

reducing the loss rate of natural habitats), using forest cover and phenology to measure the contribution of biodiversity to carbon stocks (Target 15)⁴ would ignore taxa invisible to satellites and could thus result in policymakers failing to exert the additional effort that is required to conserve saprotrophic fungal diversity, seed-dispersing mammals, and the seemingly inconsequential isopod, all of which have been implicated in boosting carbon storage^{17,33,34}. More generally, land-cover class, which is a common EO-indicator, is a highly error-prone way to map and assess the complex processes supporting ecosystem services³⁵. In short, convenient EO products could lead policymakers to focus only on that portion of biodiversity and ecosystem services that is directly observed by remote sensing, ignoring the rest.

High-Throughput Biodiversity Measurement

Most biodiversity, whether animal, fungal, plant, or microbial, and its many functions and services, is invisible to EO and will remain so for some time. But a growing number of efficient technologies are available for detecting and identifying biodiversity on the ground^{36,37} (**Supplementary Note 3** “Biodiversity technology”). Automated bioacoustic and camera-trap recording devices (ARDs) can run continuously for weeks and accumulate thousands of records of invertebrates, birds, fish, reptiles, amphibians, and

mammals, and thus allow extended sampling of large areas at low workloads³⁸⁻⁴².

Alternatively, high-throughput DNA sequencers can be used in metabarcoding or metagenomic pipelines to detect and identify anywhere from one to thousands of species at a time from mass-collected, bulk samples of organisms (e.g. 'biodiversity soups'⁴³), or from 'environmental DNA,' which is DNA liberated into the environment in the skin, hair, mucous, saliva, sperm, eggs, exudates, faeces, urine, blood, spores, root fragments, leaves, fruit, pollen, or rotting body parts of their original owners^{44,45} (**Figure 2, Supplementary Note 3**). Multiple studies have now shown that metabarcode datasets reflect high-quality, morphologically identified biodiversity datasets sufficiently closely to allow correct management decisions, given best-practice protocols and controls⁴⁶⁻⁵¹.

The taxonomic identities, phylogenetic affinities, functional genes⁵², spectral properties (of visible vegetation^{26,53,54}), and/or co-occurrence patterns⁵⁵ of the detected species can be used to parameterise process-based production functions for ecosystem services⁵⁶⁻⁵⁸ (**Figure 1**). For instance, the species identities and biomasses of wild bees identified metagenomically from bulk samples⁵⁹ could be combined with flower-use observation data⁶⁰ and detailed vegetation classification from EO to infer the availability and nature of local pollination services. Metagenomic data matched to identified species can be particularly powerful when the impacts of species loss on ecosystem function are not

243 random, evidence that has previously relied on intensive field sampling, e.g. in tropical
244 freshwater⁶¹ and marine benthic communities⁶².

245 **Statistical Modelling as the Bridge**

246 Earth Observation technology can produce large-scale, fine-resolution maps and dense
247 time series of a wide range of biophysical variables (**Supplementary Note 1 and 2**), but
248 it is difficult to translate the biophysical variables into biodiversity information. In
249 contrast, ARDs and DNA sequencing are capable of generating large amounts of
250 biodiversity information at species- or even individual-level resolution^{63,64}, but only from
251 point samples (**Supplementary Note 3**). Modern methods of statistical modelling allow
252 us to interpolate these point samples to build continuous species maps and to estimate
253 emergent metrics such as richness and dissimilarity⁶⁵⁻⁶⁸, potentially also including
254 estimates of species abundance or biomass, depending on the sampling and analytical
255 methods used (**Supplementary Note 4** "Statistical modelling").

256 The three approaches with immediate potential are *Joint Species Distribution Models*⁶⁹⁻⁷²
257 (including *Latent Variable Models*), *Community Occupancy-Detection Models*⁷³, and
258 *Generalised Dissimilarity Models*^{65,74} (**Figure 3, Supplementary Note 4**). Each approach
259 starts with a site-by-species matrix, from data that have been collected by ARDs or been
260 generated via metabarcoding or metagenomics (**Figure 2, Supplementary Note 3**), plus

any existing species distribution data. If some species are not detected, repeat sampling can be used to infer missing occurrences⁷³. The site-by-species matrix is then paired with a corresponding site-by-environmental-covariate matrix, generated from continuous EO data plus any relevant geographical layers, and the two datasets are combined statistically to infer the joint distributions of multiple species across entire regions (**Figure 3, Supplementary Note 4**). All three approaches also provide a rigorous framework for quantifying sources of uncertainty and have already been applied successfully to conventionally acquired datasets (**Box 3**).

Box 3. Current Practice in Community Modelling

Ovaskainen et al.⁷¹ used a joint species distribution model to predict the distributions of 55 butterfly species scored for presence/absence on a grid of 2609 10 X 10-km cells across Great Britain that had been sampled from 1995-1999 in a large citizen-science project. The model was successfully parameterised with a training dataset of just 300 cells and four environmental covariates (degree-days and three types of vegetation cover), plus spatially structured latent variables. Latent variables use observed species subgroupings to detect the effects of unmeasured environmental filters or species interactions such as competition. The parameterised model was used to predict butterfly

communities in the testing dataset, which consisted of the remaining 2309 grid cells.

Together, the measured and latent variables explained an average of 42% of the variance in species occurrence (with medium-prevalence species more accurately predicted), and the two most dominant latent variables revealed a north-south gradient in species composition, with especially distinct communities in the southeast and northwest.

Species richness per grid cell was accurately predicted, and the model's ability to discriminate presence and absence was high (mean AUC = 0.91).

Kéry and Royle⁷⁵ used community-occupancy modelling to analyse the 2001 Swiss breeding-bird survey while accounting for variation in detectability due to season, site, and species effects. The dataset consisted of 254 1-km² grid cells, each visited three times. The fitted model predicted each species' probability of occurrence as a function of site elevation and forest cover, as well as variance in the uncertainty of occurrence estimates, making it possible to estimate species distributions across the landscape and confidence in those estimates. Parameter estimates were naturally less precise for rare species, but information could be 'borrowed' from data-rich species to increase the precision of predictions for rare species. These procedures were able to compensate for the fact that only 134 total bird species had been detected in the survey, which is less than the true total of 163 species known to breed regularly in Switzerland, plus 22

occasional residents (the testing dataset). The occupancy-corrected model estimated that between 1 and 11 species had been overlooked per grid cell and thus, that the true total in 2001 was 169 species.

Mokany et al.⁷⁶ applied Generalised Dissimilarity Modelling (GDM) to a dataset of 2330 expert surveys of New Zealand land snails, which recorded 845 of 998 known species. The GDM was parameterised with a training dataset of 2280 surveys and fourteen environmental variables and explained 57% of the variation in beta diversity. In addition, a generalised additive model parameterised on the training dataset explained 27% of the variation in species richness (after scaling the 20 x 20-m survey quadrats to match the area of modelling units (200 x 200-m); see discussion of scaling in **Supplementary Note 4**). Finally, the outputs were combined using a procedure called DynamicFOAM to assign snail species to communities across New Zealand. Error was assessed by predicting compositions in a testing dataset of 50 sites that had been held out of the model. On average, the model was able to predict half the species that had been observed in each cell, and the predicted total occupancy area per species was highly correlated with the number of quadrat occurrences (Pearson's $r = 0.902$). When quadrats were pooled into groups of 3 to 400 to reduce sampling stochasticity, predicted species richnesses almost perfectly explained observed richnesses ($R^2 = 0.99$).

316

317

318 By mapping species distributions as the primary output, we do not lock ourselves into an
319 arbitrary set of convenient indicators, and ongoing discoveries on the relationship
320 between biodiversity and function, which are typically carried out at the species level, can
321 be added. As an illustration, the species diversity of wood-decaying fungi in natural
322 forests is notoriously difficult to assay but can be predicted in part by the volume and
323 species diversity of the stock of dead wood on the ground⁷⁷, and these environmental
324 covariates are partially quantifiable via airborne LiDAR sensors (**Supplementary Note**
325 **1**)⁷⁸, thus allowing EO-based inference of the distribution and level of wood-decaying
326 fungal diversity. Subsequent and unrelated research has suggested that pieces of dead
327 wood inhabited by a higher diversity of fungal species decompose more slowly, possibly
328 due to more intense interference competition³⁴. Combining the two results suggests that
329 an EO-derived map of fungal species diversity could be used to contrast landscape
330 management options for how well they conserve saprotrophic fungal biodiversity and
331 thus enhance carbon storage.

332 Two further reasons for focusing on species-resolution maps as the primary output are
333 that the regional species pool (*gamma diversity*) and the biological dissimilarity of sites

(*beta diversity*) could contribute to maintaining functional stability^{58,79,80} and that species-resolution outputs retain the option of aggregation to represent different aspects of biodiversity, including higher-taxonomic, functional, and phylogenetic groupings⁸¹. Many methods are also available to predict *individual* species ranges, and EO can help improve their accuracy, as shown by an example⁸² combining MODIS satellite data with environmental DNA to map an invasive diatom over a watershed [Target 9, invasive species pathway identified] (**Supplementary Figure 3.1**). However, ecosystem functions and services are rarely delivered by only one species, and simply summing the outputs of individual models to simulate communities is computationally inefficient, statistically flawed, and does not account for species interactions⁸³.

From CEOBE to Aichi

In essence, our argument is that new technologies make the new community-modelling approaches (**Box 3, Figure 3**) widely feasible, especially in biodiversity hotspots, where it is particularly difficult to generate large datasets. Larger numbers of environmental covariates and species together increase explanatory power by providing a greater breadth of predictors, and by exploiting latent variables and letting rare species 'borrow' information^{42,75,84}, respectively. As a result, continuous streams of EO data can be more powerfully interpreted to track biodiversity status and trends (**Figure 1**).

The predictive performance of fitted models can be cross-validated by rounds of comparison with testing datasets that were either split from the model-training dataset^{71,76} or derived from historical and expert knowledge⁷⁵, and thus, the adequacy of the input data and sampling design, or conversely the degree of model uncertainty, can be assessed *post hoc* (**Box 3**). The regularly updated biodiversity maps that are the primary outputs of the CEOBE approach (**Figure 1**), plus the quantified uncertainty in those maps, can then be incorporated into a larger process of structured decision making and adaptive management⁸⁵⁻⁸⁷ to (1) identify likely consequences of proposed actions by observing natural experiments that mimic those actions, (2) compare observed results of management interventions against objectives, and (3) help identify and tackle sources of uncertainty.

An early example of the CEOBE approach is given by Sollmann et al.⁴², who used community-occupancy modelling to connect environmental covariates from the 5-m-resolution RapidEye satellite to point-sample data from camera traps in three tropical-forest logging concessions in Sabah, Malaysian Borneo, one of which has been managed to reduced-impact-logging standards set by the Forest Stewardship Council (Aichi Target 7, sustainable management under forestry). The dataset consisted of detection events for 28 mammal species at 166 camera-trap stations, each station scored using EO data for

370 distance to water, distance to oil-palm plantation, and forest condition. Estimated
371 relationships between species occurrence and the three covariates were used to predict
372 species occurrence across the three reserves, with rare mammal species borrowing
373 information from more common ones. Species richness was estimated to be higher in
374 the FSC-certified reserve, particularly for threatened species (Target 12, improved
375 conservation status of threatened species). The percentage of area occupied, which could
376 indicate larger population sizes, was also estimated to be higher in the FSC-certified
377 reserve for the majority of species, including for some highly endangered species like the
378 Sunda pangolin *Manis javanica*. Finally, the modelled species richness maps were found
379 to correlate strongly with EO-estimated aboveground biomass at the large spatial grain
380 of whole reserves, but not at a finer resolution (potentially due to hunting at reserve
381 borders), further demonstrating the critical contribution of ground-level point samples
382 for linking pure-EO data to biodiversity.

383 The major remaining components of uncertainty relate to generalisability, because only a
384 single FSC-certified reserve was sampled; the applicability of results to arboreal species,
385 which tend to be detected more frequently in forests with disturbed canopy but are not
386 necessarily more widespread in these forests; and wide confidence intervals around
387 parameter estimates for some species as a consequence of sparse data and a fairly

388 complex hierarchical model. This example serves as a proof of concept that camera
389 trapping and occupancy modelling can be used to assess biodiversity conservation based
390 on species maps, and the approach has been incorporated in the ten-year forest
391 management plan and wildlife monitoring strategy for the FSC-certified area. Repeated
392 surveys will help to narrow uncertainties in the model, and a future power analysis is
393 planned to estimate the sampling effort required to detect trends and/or provide
394 estimates with a desired level of certainty⁸⁸.

395 Another example of the CEOBE approach is the use of Generalised Dissimilarity
396 Modelling to connect EO-derived metrics of habitat degradation and fragmentation^{89,90}
397 to over 300 million records of more than 400,000 species from the Global Biodiversity
398 Information Facility (www.gbif.org) and the Map of Life (mol.org)⁹¹. The GDM models
399 spatial turnover in biodiversity composition at 1-km-resolution globally, and by invoking
400 the assumption that terrestrial biodiversity declines according to the classical species-
401 area power function, the GDM estimates the proportion of biodiversity that has been
402 retained in each grid cell after habitat loss, based on the proportion of similar habitat
403 remaining unimpacted within the landscape⁹². This metric thus tracks whether rates of
404 loss, degradation, and fragmentation of natural habitats are being reduced (Aichi Target
405 5). Further, by combining this approach with a global database of protected-area

406 coverage (www.protectedplanet.net), it is possible to report progress against Target 11,
407 which aims for protected areas to cover areas of particular importance to biodiversity
408 and ecosystem services and to be ecologically representative and connected (see also
409 Ref. 93). An important caveat is that the biodiversity data in this case are historical in
410 nature and thus contain the taxonomic and sampling biases and constraints of the past
411 (**Box 2**). Ideally, the biodiversity data will transition to up-to-date, properly sampled, and
412 more taxonomically comprehensive point samples.

413 Of course, CEOBE outputs cannot contribute to all Aichi Targets, namely those that are
414 focused on policy, planning, and funding reform (Targets 2, 3, 4, 20), the conservation of
415 genetic cultivars (Target 13), the alleviation of climate-change pressures on coral reefs
416 (Target 10), benefits sharing (Target 16), and the integration of traditional knowledge
417 (Target 18). It also remains to be seen how well or poorly EO data reflect biodiversity in
418 aquatic ecosystems (Targets 6 and 11), although environmental DNA on its own is a
419 highly promising source of data on aquatic biodiversity. On the other hand, the efficient
420 production of biodiversity maps and open access to analytical pipelines will help to
421 disseminate the science base and technologies related to biodiversity (Target 19), and
422 could contribute to public awareness of efforts to conserve biodiversity (Target 1) and
423 improve the efficiency of national biodiversity planning (Target 17).

Conclusions

It is extremely difficult to identify all the species present in a location (*the Linnaean challenge*), to delimit the geographic distributions of species (*the Wallacean challenge*), and to quantify their responses to natural and anthropogenic environmental change (*the Hutchinsonian challenge*)⁹⁴. A synergy of Earth Observation, automated recording devices, high-throughput DNA sequencing, and modern statistical modelling can meet these challenges by making it possible to scale up from data-rich but finite sets of point samples to spatially continuous biodiversity maps, which are more informative than a few convenient indicator species but still let us generate summary statistics to communicate trends to decision-makers and the general public. The use of formal statistical frameworks lets us quantify error, identify gaps in our understanding, objectively rank the most likely pressures on biodiversity from multiple candidates, and increase the robustness of change detection. Adding information on species interactions and functions helps link biodiversity to ecosystem functions and services (**Box 1, Figure 1**) in a process-based approach⁵⁶, rather than relying on crude estimates from land classes³⁵. Finally, as DNA-based technologies mature, the same samples could track population-genetic diversity^{64,95,96}.

441 A global, multi-resolution monitoring network is thus within our reach but will still
442 involve a number of challenges associated with technical capacity, computation and data
443 storage, and data standardisation. For every ecologically distinct region, there will be an
444 initial cost to collect data for model parameterisation, followed by a low level of
445 continuous sampling, which will be necessary for updating models and for surveillance
446 monitoring of environmental drivers that are invisible to EO, such as broad-spectrum
447 insecticides. The initial costs are probably best borne by governments, as part of their
448 commitment to the Convention on Biological Diversity, and there is great promise in
449 using citizen-science networks to collect standardised, bulk biodiversity samples over
450 large areas. A laudable example is the School Malaise Trap Program that recruited
451 hundreds of secondary-school science classes to collect arthropods across Canada
452 (malaiseprogram.com). Initial investment could also come from existing monitoring
453 budgets with the expectation that additional information content will compensate for
454 reduced sample numbers within existing programs⁸². The follow-up continuous sampling
455 requires steady funding streams, and the standardisation of the CEOBE approach meets
456 the needs of international certification schemes, such as REDD+, Climate, Community &
457 Biodiversity Standards, Forest Stewardship Council, and the Roundtable on Sustainable
458 Palm Oil, which all require the continuous monitoring of biodiversity and ecosystem

459 services. Biodiversity-offset payments to mitigate the impacts of development and
460 carbon emissions are also expected to provide funding streams, and standardised
461 assessments are needed to ensure that offsetting results in biodiversity net gain⁹⁷.

462 The CEOBE approach also depends on institutional support for the multidisciplinary
463 collaborations needed to generate, combine, analyse, and act upon data from disparate
464 disciplines (EO, ARDs, genomics, taxonomy and systematics, ecosystem functions and
465 services, statistics, and decision science), expertise that no single individual has^{12,30,98}.

466 Identifying causal determinants of species distributions needs a clear understanding of
467 phylogenetic structure and functional diversity, the ecological processes involved, and
468 what EO sensors can and cannot observe⁹⁹. Expert knowledge will also contribute to
469 sampling design and covariate selection so that the full breadth of environmental
470 conditions is captured, especially those not visible to EO.

471 On the other hand, collaborations need not be global. Political and social interests will
472 vary by region, and agencies should be encouraged to trial CEOBE within their
473 jurisdictions where there are clear opportunities to improve management, while also
474 enforcing the publication of primary data and analytical pipelines^{27,100}. The
475 Intergovernmental Platform on Biodiversity and Ecosystem Services (IPBES) could play an
476 important role as a global coordinating institution.

477 Resources for environmental management are always likely to be limited, but by doing
478 more with our expensively gained field data, we can take action more efficiently and
479 effectively. What is required now is leadership by governments and international
480 organisations to stimulate integrated research and to endorse the use of comprehensive
481 biodiversity information⁶.

482 **Acknowledgements**

483 This article is a product of the EO-BESS Working Group, organised by Heiko Balzter,
484 David Raffaelli, and Beth Cole and funded by the UK Natural Environment Research
485 Council. Individual author acknowledgements are in Supplementary Information.

486 **Author Contributions**

487 BC and HB led the sections on Earth Observation technology. KB and DWY led the
488 sections on Biodiversity technology. AB led the sections on Statistical modelling. AB, RS,
489 AW, OO, and DWY led the sections on case studies (Box 3 and CEOBE to Aichi). CM led
490 the Conclusions section. Figures were created by KB, AB, CC, and AZ. All authors
491 contributed to multiple rewrites, with a large contribution by DR. AB and DWY wrote the
492 first draft and supervised the work.

493 **Additional Information**

494 **Correspondence should be addressed to DWY.**

Competing Interests

DWY and AV are co-founders of a private company that provides commercial metabarcoding services.

References

- 1 Verrelst, J. *et al.* Optical remote sensing and the retrieval of terrestrial vegetation biogeophysical properties – A review. *ISPRS Journal of Photogrammetry and Remote Sensing* **108**, 273-290, doi:[10.1016/j.isprsjprs.2015.05.005](https://doi.org/10.1016/j.isprsjprs.2015.05.005) (2015).
- 2 Wulder, M. A. *et al.* Virtual constellations for global terrestrial monitoring. *Remote Sensing of Environment* **170**, 62-76, doi:[10.1016/j.rse.2015.09.001](https://doi.org/10.1016/j.rse.2015.09.001) (2015).
- 3 Toth, C. & Józków, G. Remote sensing platforms and sensors: A survey. *ISPRS Journal of Photogrammetry and Remote Sensing* **115**, 22-36, doi:[10.1016/j.isprsjprs.2015.10.004](https://doi.org/10.1016/j.isprsjprs.2015.10.004) (2016).
- 4 O'Connor, B. *et al.* Earth observation as a tool for tracking progress towards the Aichi Biodiversity Targets. *Remote Sensing in Ecology and Conservation* **1**, 19-28, doi:10.1002/rse2.4 (2015).
- 5 Skidmore, A. K. *et al.* Environmental science: Agree on biodiversity metrics to track from space. *Nature News* **523**, 403, doi:10.1038/523403a (2015).
- 6 **Pettorelli, N. *et al.* Framing the concept of satellite remote sensing essential biodiversity variables: challenges and future directions. *Remote Sensing in Ecology and Conservation* **2**, 122-131, doi:10.1002/rse2.15 (2016).
Identifies candidate essential biodiversity variables derived directly from remote sensing and assesses their feasibility for global biomonitoring.**

- 517 7 CBD. Decision adopted by the Conference of the Parties to the Convention on
 518 Biological Diversity at its Tenth Meeting. Decision X/2. The Strategic Plan for
 519 Biodiversity 2011–2020 and the Aichi Biodiversity Targets.
 520 (UNEP/CBD/COP/DEC/X/2. 29 October 2010., 2010).

- 521 8 UNGA. Resolution adopted by the General Assembly on 25 September 2015.
 522 Transforming our world: the 2030 Agenda for Sustainable Development. United
 523 Nations General Assembly. Seventieth Session. A/RES/70/1., (2015).

- 524 9 UNFCCC. Adoption of the Paris Agreement. FCCC/CP/2015/L.9/Rev.1. (2015).

- 525 10 Tittensor, D. P. *et al.* A mid-term analysis of progress toward international biodiversity
 526 targets. *Science* **346**, 241-244, doi:10.1126/science.1257484 (2014).

- 527 11 Durance, I. *et al.* The Challenges of Linking Ecosystem Services to Biodiversity.
 528 *Advances in Ecological Research* **54**, 87-134, doi:10.1016/bs.aecr.2015.10.003
 529 (2016).

- 530 12 Pettorelli, N., Owen, H. & Duncan, C. How do we want Satellite Remote Sensing to
 531 support biodiversity conservation globally? *Methods in Ecology and Evolution* **7**, 656-
 532 665, doi:10.1111/2041-210X.12545 (2016).

- 533 13 IPBES. Decision and scoping report for the IPBES global assessment on biodiversity
 534 and ecosystem services. Decision IPBES-4/1. Work Programme of the Platform.
 535 (2016).

- 536 14 Dawson, T. P., Cutler, M. E. J. & Brown, C. The role of remote sensing in the
 537 development of SMART indicators for ecosystem services assessment. *Biodiversity*
 538 **17**, 136-148, doi:10.1080/14888386.2016.1246384 (2016).

- 539 15 Pereira, H. M. *et al.* Essential Biodiversity Variables. *Science* **339**, 277-278,
 540 doi:10.1126/science.1229931 (2013).

- 541 16 Proença, V. *et al.* Global biodiversity monitoring: From data sources to Essential
 542 Biodiversity Variables. *Biological Conservation*, doi:10.1016/j.biocon.2016.07.014
 543 (2016).
- 544 17 Peres, C. A., Emilio, T., Schietti, J., Desmoulière, S. J. M. & Levi, T. Dispersal
 545 limitation induces long-term biomass collapse in overhunted Amazonian forests.
 546 *Proceedings of the National Academy of Sciences USA* **113**, 892–897,
 547 doi:10.1073/pnas.1516525113 (2016).
- 548 18 Levi, T., Shepard Jr, G. H., Ohl-Schacherer, J., Peres, C. A. & Yu, D. W. Modelling
 549 the long-term sustainability of indigenous hunting in Manu National Park, Peru:
 550 landscape-scale management implications for Amazonia. *Journal of Applied Ecology*
 551 **46**, 804–814, doi:10.1111/j.1365-2664.2009.01661.x (2009).
- 552 19 Ferrier, S. Extracting More Value from Biodiversity Change Observations through
 553 Integrated Modeling. *BioScience* **61**, 96–97, doi:10.1525/bio.2011.61.2.2 (2011).
- 554 20 Belward, A. S. & Skøien, J. O. Who launched what, when and why; trends in global
 555 land-cover observation capacity from civilian earth observation satellites. *ISPRS*
 556 *Journal of Photogrammetry and Remote Sensing* **103**, 115–128,
 557 doi:[10.1016/j.isprsjprs.2014.03.009](https://doi.org/10.1016/j.isprsjprs.2014.03.009) (2015).
- 558 21 Roy, D. P. *et al.* Landsat-8: Science and product vision for terrestrial global change
 559 research. *Remote Sensing of Environment* **145**, 154–172,
 560 doi:[10.1016/j.rse.2014.02.001](https://doi.org/10.1016/j.rse.2014.02.001) (2014).
- 561 22 **Turner, W. *et al.* Free and open-access satellite data are key to biodiversity**
 562 **conservation. *Biological Conservation* 182, 173–176,**
 563 **doi:[10.1016/j.biocon.2014.11.048](https://doi.org/10.1016/j.biocon.2014.11.048) (2015).**
 564 **Identification of continuity, affordability, and accessibility as having major**

- 565 **impact on the utility of remote sensing data for tracking and understanding**
566 **biodiversity change.**
- 567 23 Butler, D. Earth observation enters next phase. *Nature* **508**, 160-161,
568 doi:10.1038/508160a (2014).
- 569 24 Berger, M., Moreno, J., Johannessen, J. A., Levelt, P. F. & Hanssen, R. F. ESA's
570 sentinel missions in support of Earth system science. *Remote Sensing of*
571 *Environment* **120**, 84-90, doi:[10.1016/j.rse.2011.07.023](https://doi.org/10.1016/j.rse.2011.07.023) (2012).
- 572 25 Malenovsky, Z. *et al.* Sentinels for science: Potential of Sentinel-1, -2, and -3
573 missions for scientific observations of ocean, cryosphere, and land. *Remote Sensing*
574 *of Environment* **120**, 91-101, doi:[10.1016/j.rse.2011.09.026](https://doi.org/10.1016/j.rse.2011.09.026) (2012).
- 575 26 **Asner, G. P. *et al.* Airborne laser-guided imaging spectroscopy to map forest**
576 **trait diversity and guide conservation. *Science* 355, 385-389,**
577 **doi:10.1126/science.aaj1987 (2017).**
- 578 **Large-scale mapping of multivariate forest canopy functional traits from**
579 **airborne laser-guided imaging spectroscopy.**
- 580 27 Petrou, Z. I., Manakos, I. & Stathaki, T. Remote sensing for biodiversity monitoring: a
581 review of methods for biodiversity indicator extraction and assessment of progress
582 towards international targets. *Biodiversity and Conservation* **24**, 2333-2363,
583 doi:10.1007/s10531-015-0947-z (2015).
- 584 28 Wulder, M. A., Masek, J. G., Cohen, W. B., Loveland, T. R. & Woodcock, C. E.
585 Opening the archive: How free data has enabled the science and monitoring promise
586 of Landsat. *Remote Sensing of Environment* **122**, 2-10,
587 doi:[10.1016/j.rse.2012.01.010](https://doi.org/10.1016/j.rse.2012.01.010) (2012).

- 588 29 Lindenmayer, D. B. & Likens, G. E. Direct Measurement Versus Surrogate Indicator
589 Species for Evaluating Environmental Change and Biodiversity Loss. *Ecosystems* **14**,
590 47-59, doi:10.1007/s10021-010-9394-6 (2011).
- 591 30 Mueller, M. & Geist, J. Conceptual guidelines for the implementation of the
592 ecosystem approach in biodiversity monitoring. *Ecosphere* **7**, e01305,
593 doi:10.1002/ecs2.1305 (2016).
- 594 31 Newton, A. C. Implications of Goodhart's Law for monitoring global biodiversity loss.
595 *Conservation Letters* **4**, 264-268, doi:10.1111/j.1755-263X.2011.00167.x (2011).
- 596 32 Smaldino, P. E. & McElreath, R. The natural selection of bad science. *Royal Society*
597 *Open Science* **3**, doi:10.1098/rsos.160384 (2016).
- 598 33 Crowther, T. W. *et al.* Biotic interactions mediate soil microbial feedbacks to climate
599 change. *Proceedings of the National Academy of Sciences* **112**, 7033-7038,
600 doi:10.1073/pnas.1502956112 (2015).
- 601 34 Yang, C. Y. *et al.* Higher fungal diversity is correlated with lower CO₂ emissions from
602 dead wood in a natural forest. *Scientific Reports* **6**, doi:10.1038/srep31066 (2016).
- 603 35 Eigenbrod, F. *et al.* The impact of proxy-based methods on mapping the distribution
604 of ecosystem services. *Journal of Applied Ecology* **47**, 377-385, doi:10.1111/j.1365-
605 2664.2010.01777.x (2010).
- 606 36 Snaddon, J., Petrokofsky, G., Jepson, P. & Willis, K. J. Biodiversity technologies:
607 tools as change agents. *Biology Letters* **9**, 20121029, doi:10.1098/rsbl.2012.1029
608 (2013).
- 609 37 Turner, W. Sensing biodiversity. *Science* **346**, 301-302,
610 doi:10.1126/science.1256014 (2014).

- 611 38 Acevedo, M. A. & Villanueva-Rivera, L. J. Using automated digital recording systems
612 as effective tools for the monitoring of birds and amphibians. *Wildlife Society Bulletin*
613 **34**, 211-214, doi:10.2193/0091-7648(2006)34[211:UADRSA]2.0.CO;2 (2006).
- 614 39 Lammers, M. O., Brainard, R. E., Au, W. W. L., Mooney, T. A. & Wong, K. B. An
615 ecological acoustic recorder (EAR) for long-term monitoring of biological and
616 anthropogenic sounds on coral reefs and other marine habitats. *The Journal of the*
617 *Acoustical Society of America* **123**, 1720-1728, doi:[10.1121/1.2836780](https://doi.org/10.1121/1.2836780) (2008).
- 618 40 Jung, K. & Kalko, E. K. V. Adaptability and vulnerability of high flying Neotropical
619 aerial insectivorous bats to urbanization. *Diversity and Distributions* **17**, 262-274,
620 doi:10.1111/j.1472-4642.2010.00738.x (2011).
- 621 41 Aide, T. M. *et al.* Real-time bioacoustics monitoring and automated species
622 identification. *PeerJ* **1**, e103, doi:10.7717/peerj.103 (2013).
- 623 42 Sollmann, R. *et al.* Quantifying mammal biodiversity co-benefits in certified tropical
624 forests. *Diversity and Distributions* **23**, 317-328, doi:10.1111/ddi.12530 (2017).
- 625 43 Yu, D. W. *et al.* Biodiversity soup: metabarcoding of arthropods for rapid biodiversity
626 assessment and biomonitoring. *Methods in Ecology and Evolution* **3**, 613-623,
627 doi:10.1111/j.2041-210X.2012.00198.x (2012).
- 628 44 Taberlet, P., Coissac, E., Hajibabaei, M. & Rieseberg, L. H. Environmental DNA.
629 *Molecular Ecology* **21**, 1789-1793, doi:10.1111/j.1365-294X.2012.05542.x (2012).
- 630 45 Bohmann, K. *et al.* Environmental DNA for wildlife biology and biodiversity
631 monitoring. *Trends in Ecology & Evolution* **29**, 358-367,
632 doi:[10.1016/j.tree.2014.04.003](https://doi.org/10.1016/j.tree.2014.04.003) (2014).
- 633 46 Ji, Y. *et al.* **Reliable, verifiable and efficient monitoring of biodiversity via**
634 **metabarcoding. *Ecology Letters* 16, 1245-1257, doi:10.1111/ele.12162 (2013).**

- 635 **Demonstrated that metabarcoding data can be as reliable as high-quality**
 636 **morphological datasets for environmental management decisions.**
- 637 47 Lejzerowicz, F. *et al.* High-throughput sequencing and morphology perform equally
 638 well for benthic monitoring of marine ecosystems. *Scientific Reports* **5**, 13932,
 639 doi:10.1038/srep13932 (2015).
- 640 48 Edwards, D. P. *et al.* Selective-logging and oil palm: multitaxon impacts, biodiversity
 641 indicators, and trade-offs for conservation planning. *Ecological Applications* **24**, 2029-
 642 2049, doi:10.1890/14-0010.1 (2014).
- 643 49 Chariton, A. A. *et al.* Emergent technologies and analytical approaches for
 644 understanding the effects of multiple stressors in aquatic environments. *Marine and*
 645 *Freshwater Research* (2015).
- 646 50 Aylagas, E., Borja, Á., Irigoien, X. & Rodríguez-Ezpeleta, N. Benchmarking DNA
 647 Metabarcoding for Biodiversity-Based Monitoring and Assessment. *Frontiers in*
 648 *Marine Science* **3**, doi:10.3389/fmars.2016.00096 (2016).
- 649 51 Visco, J. A. *et al.* Environmental Monitoring: Inferring the Diatom Index from Next-
 650 Generation Sequencing Data. *Environmental Science & Technology* **49**, 7597-7605,
 651 doi:10.1021/es506158m (2015).
- 652 52 Xue, K. *et al.* Tundra soil carbon is vulnerable to rapid microbial decomposition under
 653 climate warming. *Nature Clim. Change* **6**, 595-600, doi:10.1038/nclimate2940 (2016).
- 654 53 Asner, G. P., Knapp, D. E., Anderson, C. B., Martin, R. E. & Vaughn, N. Large-scale
 655 climatic and geophysical controls on the leaf economics spectrum. *Proceedings of*
 656 *the National Academy of Sciences* **113**, E4043–E4051,
 657 doi:10.1073/pnas.1604863113 (2016).

- 658 54 Fisher, J. B., Sweeney, S. & Brzostek, E. R. Tree–mycorrhizal associations detected
659 remotely from canopy spectral properties. *Global Change Biology* **22**, 2596-2607,
660 doi:10.1111/gcb.13264 (2016).
- 661 55 **Bohan, D. A. *et al.* Next-Generation Global Biomonitoring: Large-scale,**
662 **Automated Reconstruction of Ecological Networks. *Trends in Ecology &***
663 ***Evolution*, doi:10.1016/j.tree.2017.03.001 (2017).**
- 664 **Proposal to infer interaction networks and species functions from co-**
665 **occurrence data generated by high-throughput biodiversity methods.**
- 666 56 Barnes, A. D. *et al.* Species richness and biomass explain spatial turnover in
667 ecosystem functioning across tropical and temperate ecosystems. *Philosophical*
668 *Transactions of the Royal Society B: Biological Sciences* **37**,
669 doi:10.1098/rstb.2015.0279 (2016).
- 670 57 Brose, U. & Hillebrand, H. Biodiversity and ecosystem functioning in dynamic
671 landscapes. *Philosophical Transactions of the Royal Society B: Biological Sciences*
672 **371**, 20150267, doi:10.1098/rstb.2015.0267 (2016).
- 673 58 Burley, H. M., Mokany, K., Ferrier, S. & Laffan, S. W. Macroecological scale effects
674 of biodiversity on ecosystem functions under environmental change. *Ecol. Evol.* **6**,
675 2579-2593, doi:10.1002/ece3.2036 (2016).
- 676 59 Tang, M. *et al.* High-throughput monitoring of wild bee diversity and abundance via
677 mitogenomics. *Methods in Ecology and Evolution* **6**, 1034-1043, doi:10.1111/2041-
678 210X.12416 (2015).
- 679 60 Wood, T. J., Holland, J. M. & Goulson, D. Providing foraging resources for solitary
680 bees on farmland: current schemes for pollinators benefit a limited suite of species.
681 *Journal of Applied Ecology* **54**, 323-333, doi:10.1111/1365-2664.12718 (2017).

- 682 61 McIntyre, P. B., Jones, L. E., Flecker, A. S. & Vanni, M. J. Fish extinctions alter
683 nutrient recycling in tropical freshwaters. *Proceedings of the National Academy of*
684 *Sciences* **104**, 4461-4466, doi:10.1073/pnas.0608148104 (2007).
- 685 62 Solan, M. *et al.* Extinction and Ecosystem Function in the Marine Benthos. *Science*
686 **306**, 1177-1180, doi:10.1126/science.1103960 (2004).
- 687 63 Sunarto, Sollmann, R., A., M. & Kelly, M. J. Camera trapping for the study and
688 conservation of tropical carnivores. *Raffles Bulletin of Zoology* **28**, 21-42 (2013).
- 689 64 Sigsgaard, E. E. *et al.* Population characteristics of a large whale shark aggregation
690 inferred from seawater environmental DNA. *Nature Ecology & Evolution* **1**, 0004,
691 doi:10.1038/s41559-016-0004 (2016).
- 692 65 Ferrier, S. Mapping Spatial Pattern in Biodiversity for Regional Conservation
693 Planning: Where to from Here? *Systematic Biology* **51**, 331-363 (2002).
- 694 66 Ferrier, S. & Guisan, A. Spatial modelling of biodiversity at the community level.
695 *Journal of Applied Ecology* **43**, 393-404, doi:10.1111/j.1365-2664.2006.01149.x
696 (2006).
- 697 67 Honrado, J. P., Pereira, H. M. & Guisan, A. Fostering integration between biodiversity
698 monitoring and modelling. *Journal of Applied Ecology* **53**, 1299-1304,
699 doi:10.1111/1365-2664.12777 (2016).
- 700 68 D'Amen, M., Rahbek, C., Zimmermann, N. E. & Guisan, A. Spatial predictions at the
701 community level: from current approaches to future frameworks. *Biological Reviews*
702 **92**, 169-187, doi:10.1111/brev.12222 (2017).
- 703 69 Warton, D. I. *et al.* So Many Variables: Joint Modeling in Community Ecology. *Trends*
704 *in Ecology & Evolution* **30**, 766-779, doi:10.1016/j.tree.2015.09.007 (2015).

- 705 70 Ovaskainen, O., Abrego, N., Halme, P. & Dunson, D. Using latent variable models to
 706 identify large networks of species-to-species associations at different spatial scales.
 707 *Methods in Ecology and Evolution* **7**, 549-555, doi:10.1111/2041-210X.12501 (2016).
- 708 71 Ovaskainen, O., Roy, D. B., Fox, R. & Anderson, B. J. Uncovering hidden spatial
 709 structure in species communities with spatially explicit joint species distribution
 710 models. *Methods in Ecology and Evolution* **7**, 428-436, doi:10.1111/2041-
 711 210X.12502 (2016).
- 712 72 **Ovaskainen, O. *et al.* How to make more out of community data? A conceptual**
 713 **framework and its implementation as models and software. *Ecology Letters*,**
 714 **doi:doi:10.1111/ele.12757 (2017).**
- 715 **Presents a general joint species distribution modelling framework for**
 716 **examining how the link from environmental covariates to species occurrence**
 717 **depends on species traits and phylogenetic relationships.**
- 718 73 Dorazio, R. M. & Royle, J. A. Estimating size and composition of biological
 719 communities by modeling the occurrence of species. *Journal of the American*
 720 *Statistical Association* **100**, 389-398 (2005).
- 721 74 Ferrier, S., Manion, G., Elith, J. & Richardson, K. Using generalized dissimilarity
 722 modelling to analyse and predict patterns of beta diversity in regional biodiversity
 723 assessment. *Diversity and Distributions* **13**, 252-264, doi:10.1111/j.1472-
 724 4642.2007.00341.x (2007).
- 725 75 Kery, M. & Royle, A. J. in *Modeling demographic processes in marked populations.*
 726 *Environmental and ecological statistics, Vol. 3.* (eds D. L. Thomson, E. G. Cooch, &
 727 M. J. Conroy) (Springer, 2009).

- 728 76 Mokany, K., Harwood, T., Overton, J., Barker, G. & Ferrier, S. Combining α - and β -
 729 diversity models to fill gaps in our knowledge of biodiversity. *Ecology letters* **14**,
 730 1043-1051, doi:10.1111/j.1461-0248.2011.01675.x (2011).
- 731 77 Hottola, J., Ovaskainen, O. & Hanski, I. A unified measure of the number, volume
 732 and diversity of dead trees and the response of fungal communities. *Journal of*
 733 *Ecology* **97**, 1320-1328, doi:10.1111/j.1365-2745.2009.01583.x (2009).
- 734 78 Mücke, W., Deák, B., Schroiff, A., Hollaus, M. & Pfeifer, N. Detection of fallen trees in
 735 forested areas using small footprint airborne laser scanning data. *Canadian Journal*
 736 *of Remote Sensing* **39**, S32-S40, doi:10.5589/m13-013 (2013).
- 737 79 Pasari, J. R., Levi, T., Zavaleta, E. S. & Tilman, D. Several scales of biodiversity
 738 affect ecosystem multifunctionality. *Proceedings of the National Academy of*
 739 *Sciences* **110**, 10219-10222, doi:10.1073/pnas.1220333110 (2013).
- 740 80 Wang, S. & Loreau, M. Ecosystem stability in space: α , β and γ variability.
 741 *Ecology letters* **17**, 891-901, doi:10.1111/ele.12292 (2014).
- 742 81 Cardinale, B. J., Duffy, J. E., Gonzalez, A. & Hooper, D. U. Biodiversity loss and its
 743 impact on humanity. *Nature* **486**, 59–67, doi:10.1038/nature11148 (2012).
- 744 82 Olson, J. R., Hawkins, C. P., Mock, K., Huntington, J. & Susfalk., R. System for
 745 Mapping And Predicting Species Of Concern (SMAP-SOC), Phase I Final Report and
 746 Phase II Plan., (NASA Earth Science Division/Applied Sciences Program,
 747 Washington D.C., 2014).
- 748 83 Calabrese, J. M., Certain, G., Kraan, C. & Dormann, C. F. Stacking species
 749 distribution models and adjusting bias by linking them to macroecological models.
 750 *Global Ecology and Biogeography* **23**, 99-112, doi:10.1111/geb.12102 (2014).

- 751 84 Ovaskainen, O. & Soininen, J. Making more out of sparse data: hierarchical modeling
752 of species communities. *Ecology* **92**, 289-295, doi:10.1890/10-1251.1 (2011).
- 753 85 Ferretti, V. & Pomarico, S. Ecological land suitability analysis through spatial
754 indicators: An application of the Analytic Network Process technique and Ordered
755 Weighted Average approach. *Ecological Indicators* **34**, 507-519,
756 doi:[10.1016/j.ecolind.2013.06.005](https://doi.org/10.1016/j.ecolind.2013.06.005) (2013).
- 757 86 Marcot, B. G. *et al.* Recent advances in applying decision science to managing
758 national forests. *Forest Ecology and Management* **285**, 123-132,
759 doi:[10.1016/j.foreco.2012.08.024](https://doi.org/10.1016/j.foreco.2012.08.024) (2012).
- 760 87 Gregory, R., Long, G., Colligan, M., Geiger, J. G. & Laser, M. When experts disagree
761 (and better science won't help much): Using structured deliberations to support
762 endangered species recovery planning. *Journal of Environmental Management* **105**,
763 30-43, doi:[10.1016/j.jenvman.2012.03.001](https://doi.org/10.1016/j.jenvman.2012.03.001) (2012).
- 764 88 Steidl, R. J., Hayes, J. P. & Schaubert, E. Statistical power analysis in wildlife
765 research. *Journal of Wildlife Management* **61**, 270-279 (1997).
- 766 89 Hansen, M. C. *et al.* High-Resolution Global Maps of 21st-Century Forest Cover
767 Change. *Science* **342**, 850-853, doi:10.1126/science.1244693 (2013).
- 768 90 Newbold, T. *et al.* Has land use pushed terrestrial biodiversity beyond the planetary
769 boundary? A global assessment. *Science* **353**, 288-291,
770 doi:10.1126/science.aaf2201 (2016).
- 771 91 GEO BON. (ed Group on Earth Observations Biodiversity Observation Network
772 Secretariat) (Leipzig, 2016).

- 773 92 Allnutt, T. F. *et al.* A method for quantifying biodiversity loss and its application to a
774 50-year record of deforestation across Madagascar. *Conservation Letters* **1**, 173-
775 181, doi:10.1111/j.1755-263X.2008.00027.x (2008).
- 776 93 Ferrier, S. *et al.* Mapping more of terrestrial biodiversity for global conservation
777 assessment. *BioScience* **54**, 1101-1109 (2004).
- 778 94 Cardoso, P., Erwin, T. L., Borges, P. A. V. & New, T. R. The seven impediments in
779 invertebrate conservation and how to overcome them. *Biological Conservation* **144**,
780 2647-2655 (2011).
- 781 95 Fitzpatrick, M. C. & Keller, S. R. Ecological genomics meets community-level
782 modelling of biodiversity: mapping the genomic landscape of current and future
783 environmental adaptation. *Ecology Letters* **18**, 1-16, doi:10.1111/ele.12376 (2015).
- 784 96 Crampton-Platt, A., Yu, D. W., Zhou, X. & Vogler, A. P. Mitochondrial metagenomics:
785 letting the genes out of the bottle. *GigaScience* **5**, 1-11, doi:10.1186/s13742-016-
786 0120-y (2016).
- 787 97 Maron, M., Gordon, A., Mackey, B., Possingham, H. P. & Watson, J. E. M. Stop
788 misuse of biodiversity offsets. *Nature* **523**, 401–403 (2015).
- 789 98 Palumbo, I. *et al.* Building capacity in remote sensing for conservation: present and
790 future challenges. *Remote Sensing in Ecology and Conservation* **3**, 21-29,
791 doi:10.1002/rse2.31 (2016).
- 792 99 Dafforn, K. A., Johnston, E. L. & Ferguson, A. Big data opportunities and challenges
793 for assessing multiple stressors across scales in aquatic ecosystems. *Marine and*
794 *Freshwater Research* **67**, 393-413 (2015).
- 795 100 Schmeller, D. S. *et al.* Towards a global terrestrial species monitoring program.
796 *Journal for Nature Conservation* **25**, 51-57, doi:[10.1016/j.jnc.2015.03.003](https://doi.org/10.1016/j.jnc.2015.03.003) (2015).

Figure legends

Figure 1. CEOBE – Connecting Earth Observation to Biodiversity and Ecosystems. Top

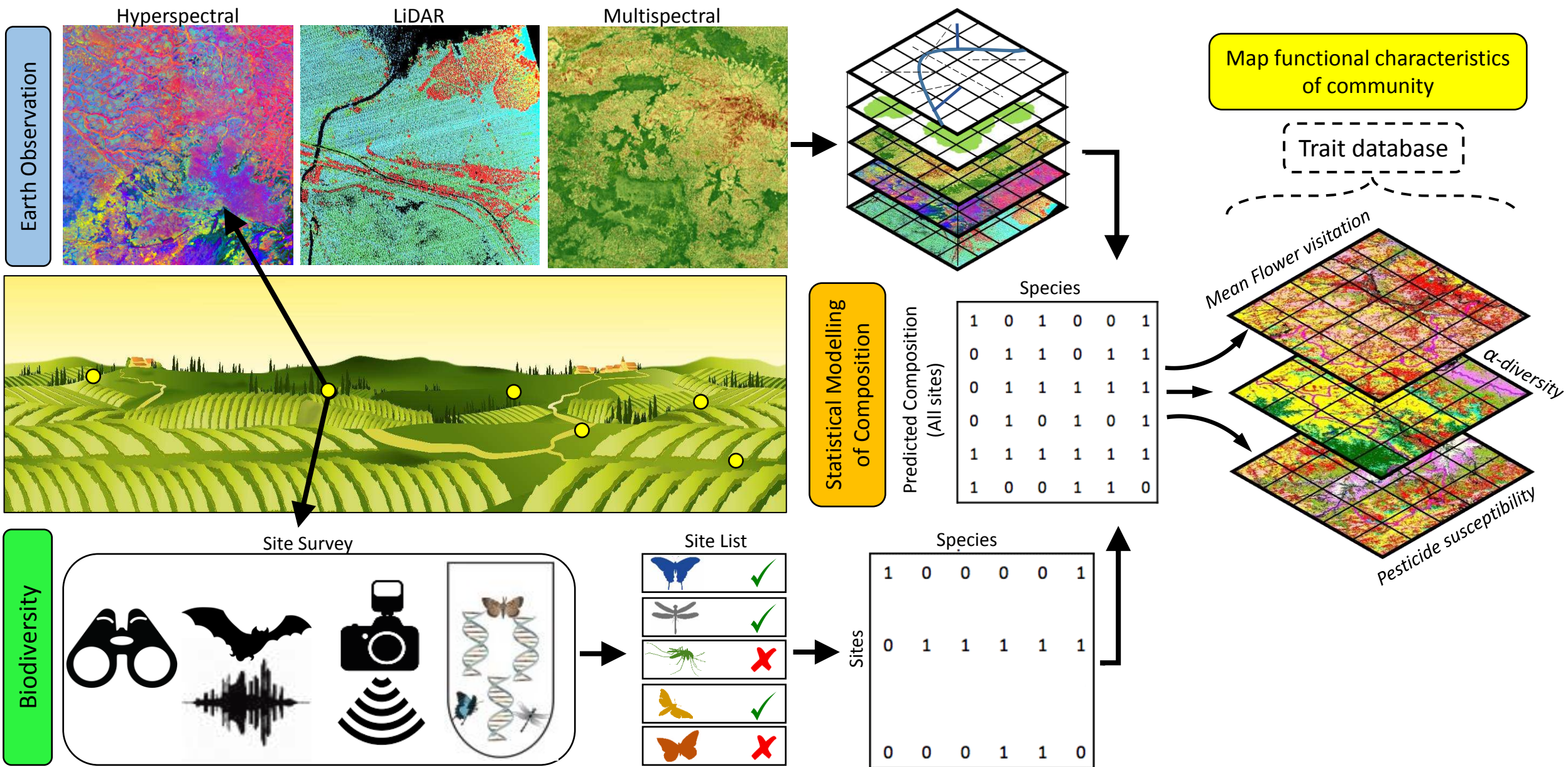
row left: EO data and other geographical datasets are used to generate spatially continuous maps of biophysical data (**S1, S2**). **Middle row left:** A real landscape with point-sample locations indicated by yellow dots. **Bottom row left:** Biodiversity is recorded manually using traditional methods, automated audio or image recording devices, or metabarcoding or metagenomic pipelines to generate a site X species table (**Figure 2, S3**). However, most of the landscape is not sampled (empty rows in the table). **Right side:** The point samples are combined statistically with continuous biophysical maps to predict biodiversity composition over the whole landscape (**S4**). In combination with ancillary data like trait databases, process-based models can then identify the functional composition of any location and map the expected distributions of ecosystem functions and services.

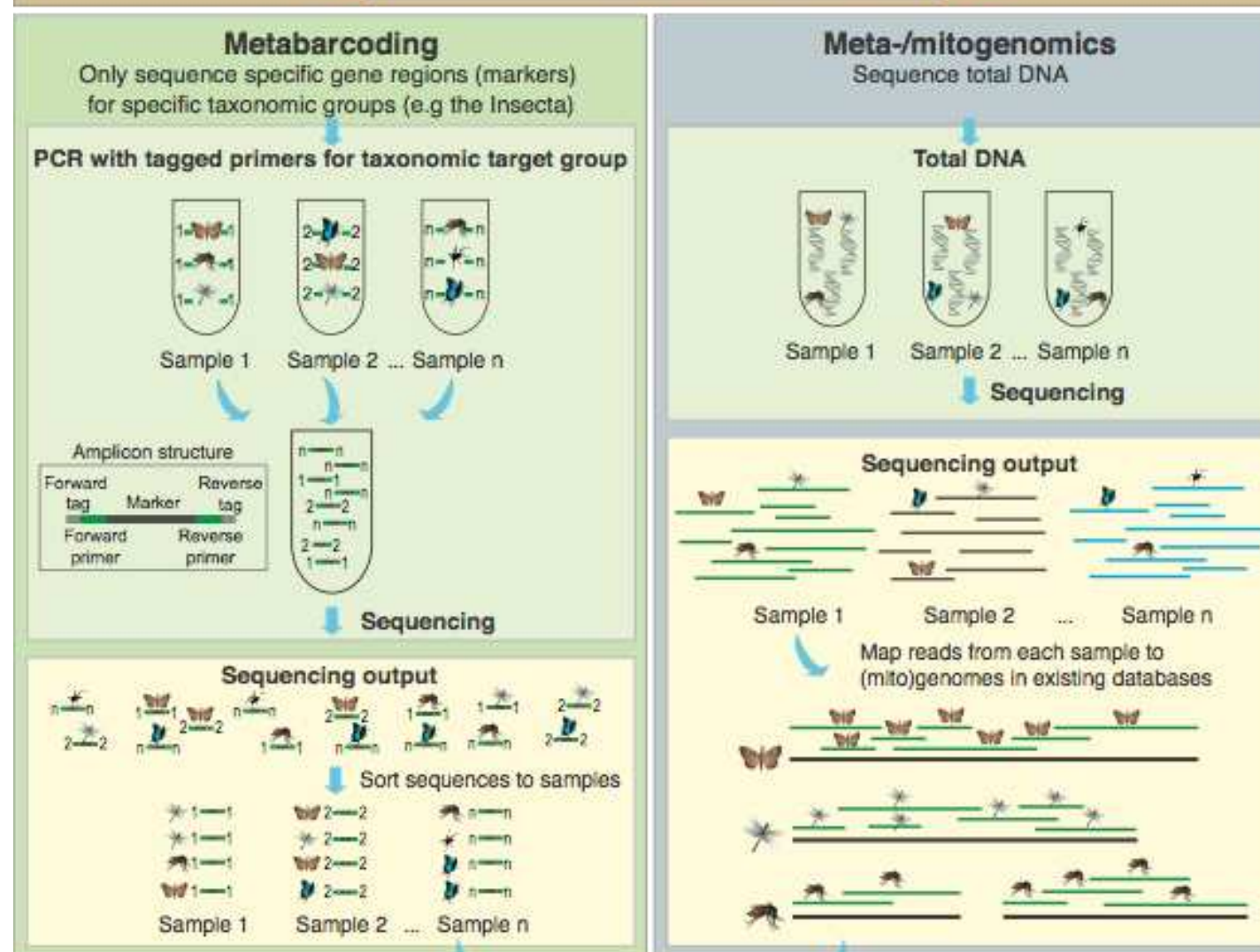
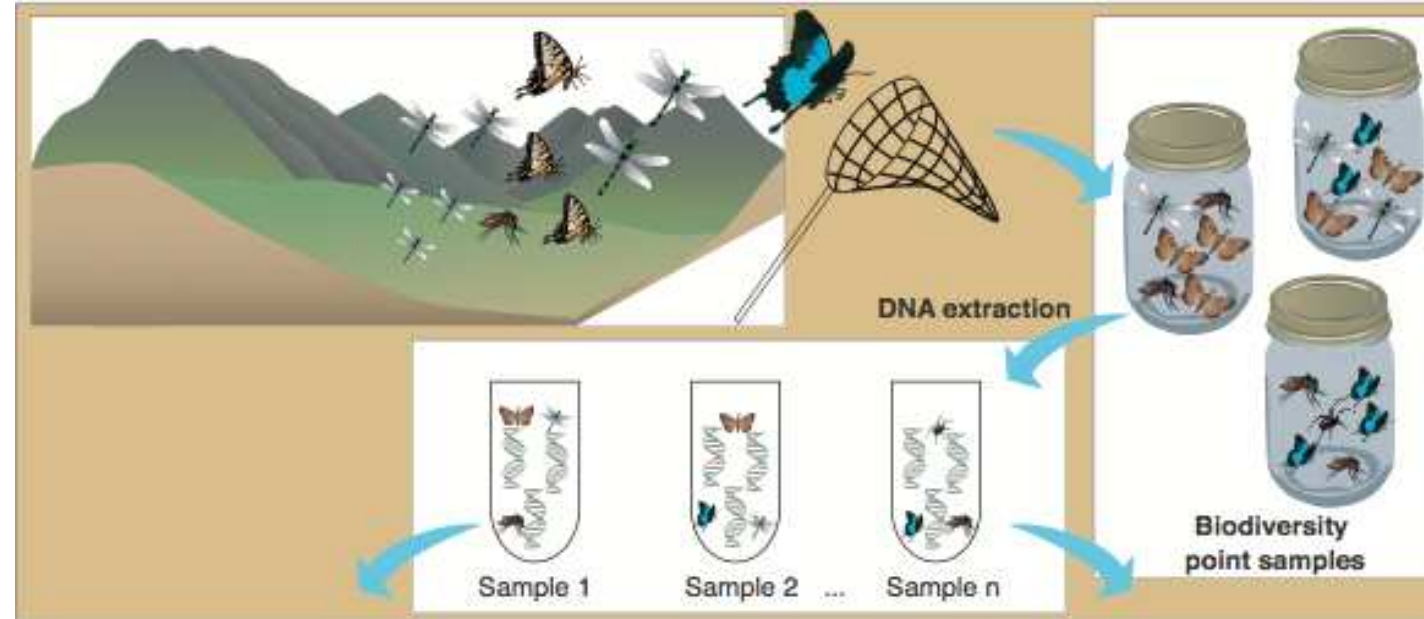
Figure 2. Metabarcoding and metagenomic processing pipelines for high-throughput

biodiversity surveys. Top row: Point locations across a landscape are sampled for biodiversity, and DNA is separately extracted from each sample. Three common sample types are (i) bulk samples of arthropods (depicted here), (ii) environmental DNA (eDNA)

815 from soil, water, and air, and (iii) invertebrate collectors of vertebrate DNA (iDNA), such
816 as mosquitoes, leeches, flies, dung beetles, and ticks. **Left column:** Metabarcoding –
817 Each sample's DNA is amplified via PCR (polymerase chain reaction) for a particular
818 marker gene that is taxonomically informative, the samples are pooled and sequenced
819 on a high-throughput sequencer, and then sorted back to sample by the sample-specific
820 tags added during PCR. The sequences are then clustered into Operational Taxonomic
821 Units (OTUs), which are species hypotheses, and assigned taxonomies by matching
822 against online databases. **Right column:** Meta/mitogenomics – Each sample's total DNA
823 is sequenced, and the output DNA reads are matched to reference genomes, which are
824 often mitochondrial genomes. **Bottom row:** The output of both processing pipelines is a
825 'sample X species' table. Metabarcoding pipelines are useful for general biodiversity
826 discovery and surveys because online barcode databases are more taxonomically
827 complete, and even without taxonomic assignment, it is possible to calculate community
828 metrics from OTUs only. Metagenomic pipelines are more costly, but advantageous when
829 it is important to reliably identify particular sets of species and to a greater extent
830 preserve relative biomass information. See **S3** for further details. Clip-art courtesy of the
831 Integration and Application Network, University of Maryland Center for Environmental
832 Science (ian.umces.edu/symbols/).

833 **Figure 3. Three statistical pathways to map community composition and summary**
834 **metrics from the combination of biodiversity point samples and continuous Earth**
835 **Observation (EO) maps.** Local diversity – α , species turnover – β , and regional diversity –
836 γ . For clarity, the figure only considers models for species occurrence (OCC), not
837 abundance. GAM: Generalised Additive Model. DynamicFOAM is described in Ref. 76.
838 See **S4** for further details.





Metabarcoding is a targeted and cost-effective approach in which only short marker(s) for the taxonomic groups desired for a given biodiversity assessment are sequenced. It is more likely to detect low-biomass taxa than is mito-/metagenomics. Metabarcoding exploits existing reference databases, which are larger than reference database collections for whole (mito)genomes.

		Species									
Sample	0	1	0	1	0	0	1	0	1	1	0
	1	0	0	0	0	1	0	0	0	0	0
	1	0	1	0	1	1	1	1	0	1	1
	0	1	0	0	0	0	0	0	1	0	0
	1	1	0	1	1	1	1	1	1	0	1
	0	0	0	1	0	1	0	1	1	0	0
	1	0	0	0	0	1	1	0	0	0	0
	0	0	0	0	0	0	0	0	0	0	0

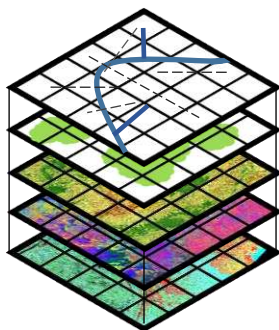
Meta-/mitogenomics requires deeper sequencing than metabarcoding because total DNA is sequenced, and only a small fraction of the sequencing output is used for detecting species. Meta-/mitogenomics relies on whole (mito-)genome reference databases, but when these are available, it has higher certainty of taxonomic assignment than does metabarcoding.

Joint Species Distribution Models / Latent Variable Models

Biodiversity point samples

EO Spatial covariates

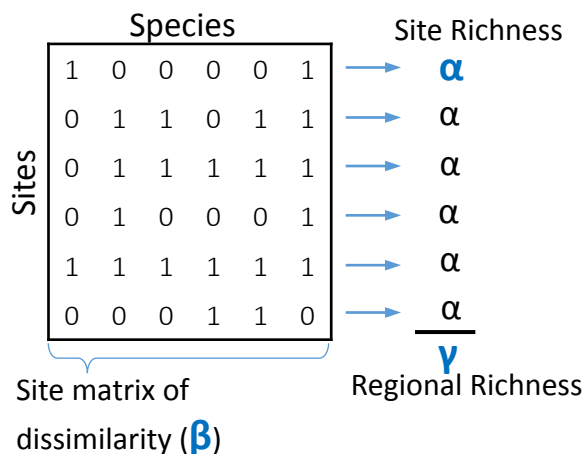
		Species					
Sites	1	1	0	0	0	0	1
	2	0	1	1	1	1	1
	3	0	1	0	0	0	1
	4	0	0	0	1	1	0
	5	0	0	0	1	1	0



$$OCC = f(\text{Site covariates}) + f(\text{Latent Variables})$$

Species distributions are described as a function of unobserved latent factors as well as observed covariates. Account for species covariance, but do not easily account for differences in species detection.

Predicted probabilities of species occurrences at all sites

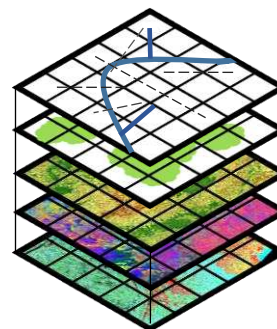


Occupancy-Detection Models

Biodiversity point samples with repeated surveys

EO Spatial covariates

		Species					
Sites	1	1	0	0	0	0	1
	2	0	1	1	1	1	1
	3	0	1	0	0	0	1
	4	0	0	0	1	1	0
	5	0	0	0	1	1	0



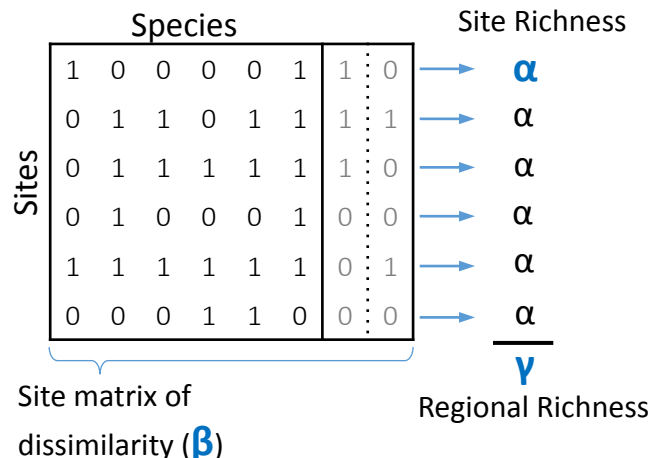
Site covariates

Survey covariates

$$OCC = \text{Occupancy.model} * \text{Detection.model}$$

Environmental covariates can describe both a species' distribution and how that distribution is observed, which itself can depend upon survey characteristics. Account for imperfect detection, but treat species independently.

Predicted probability of species occurrence at all sites (including unobserved species)

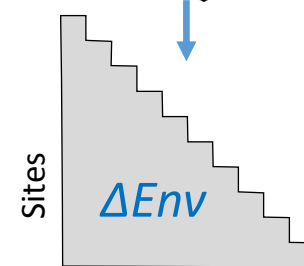
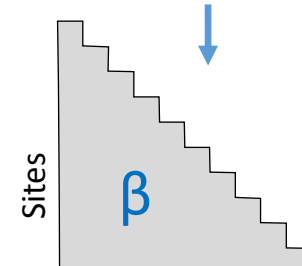
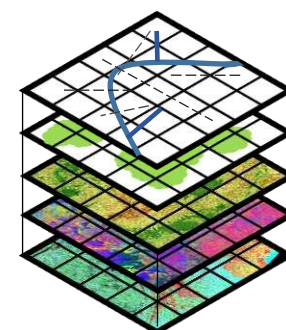


Generalised Dissimilarity Models

Biodiversity point samples (high species diversity)

EO Spatial covariates

		Species									
Sites	1	1	0	0	0	0	1	0	0	1	0
	2	1	1	0	1	1	1	0	1	0	0
	3	0	0	0	0	1	1	0	0	0	1
	4	0	1	0	0	0	0	1	0	1	0
	5	1	1	0	0	0	1	0	0	0	0



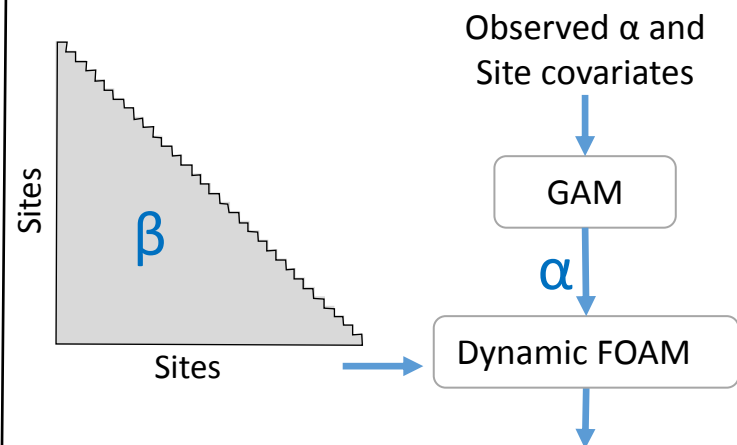
Sites

Sites

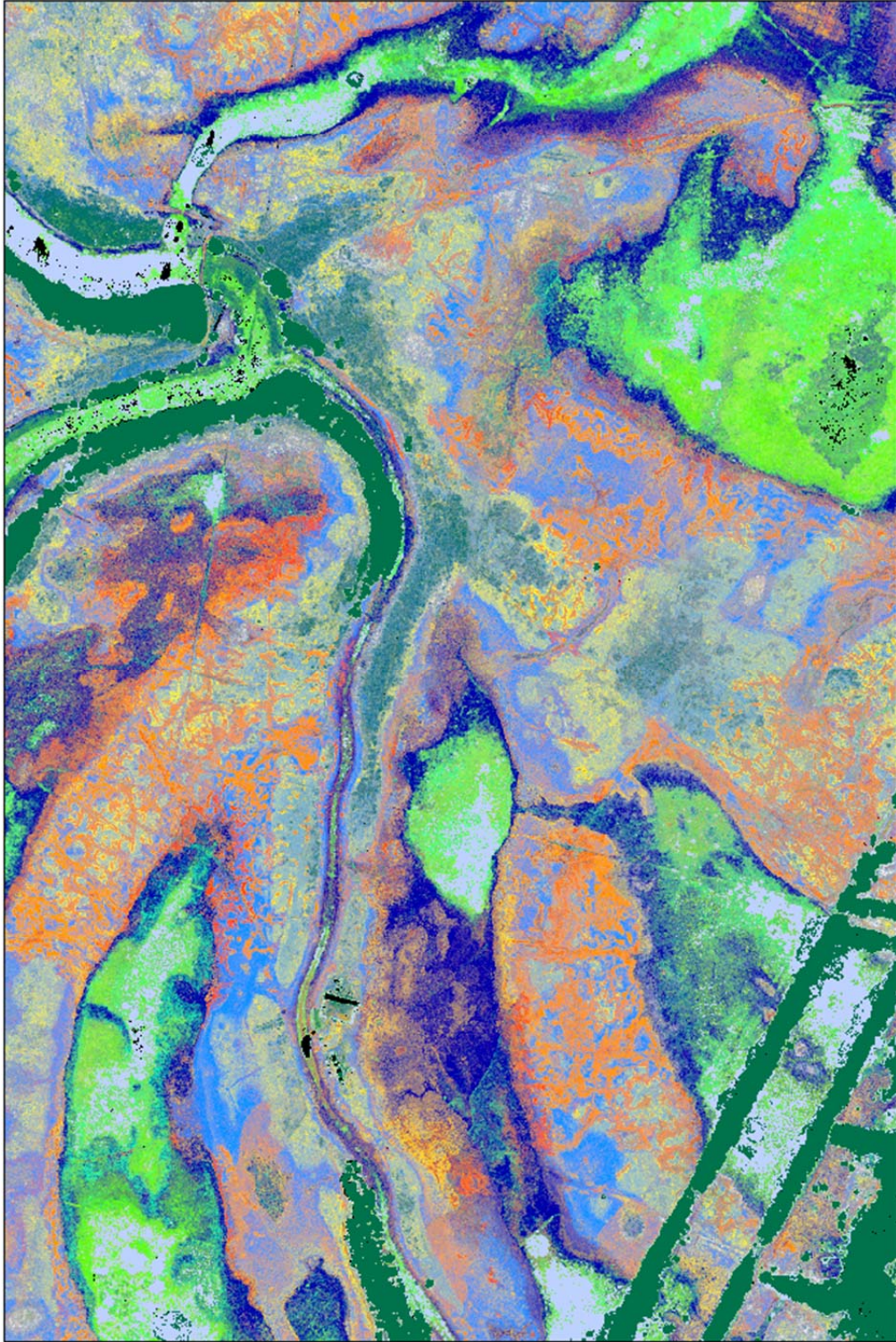
$$\beta_{ij} = f(|Env_i - Env_j|)$$

Compositional dissimilarity (β) between each pair of sites (i and j) is a function of the difference in environmental conditions (ΔEnv).

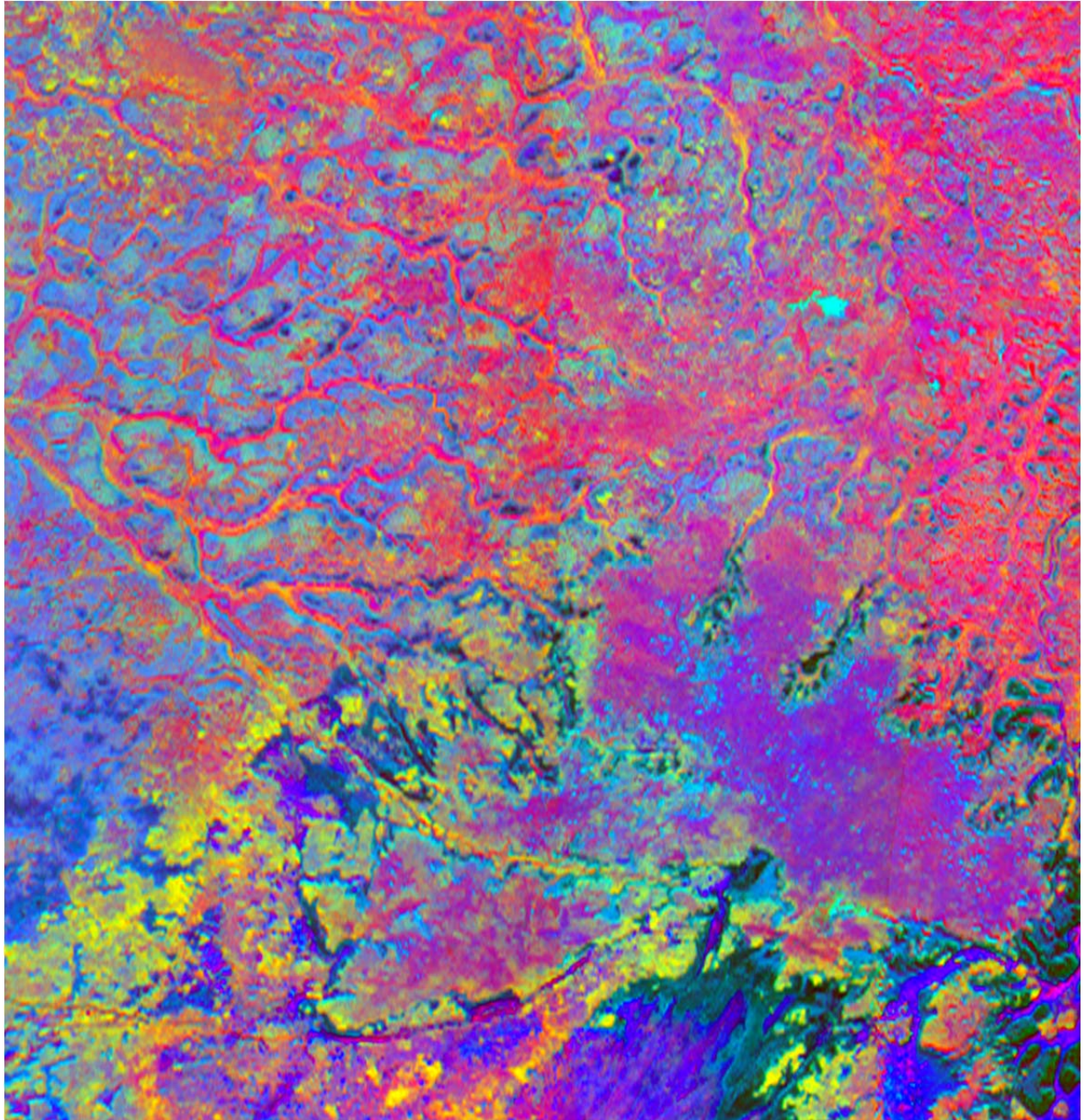
Predicted compositional dissimilarity between any pair of sites (β)



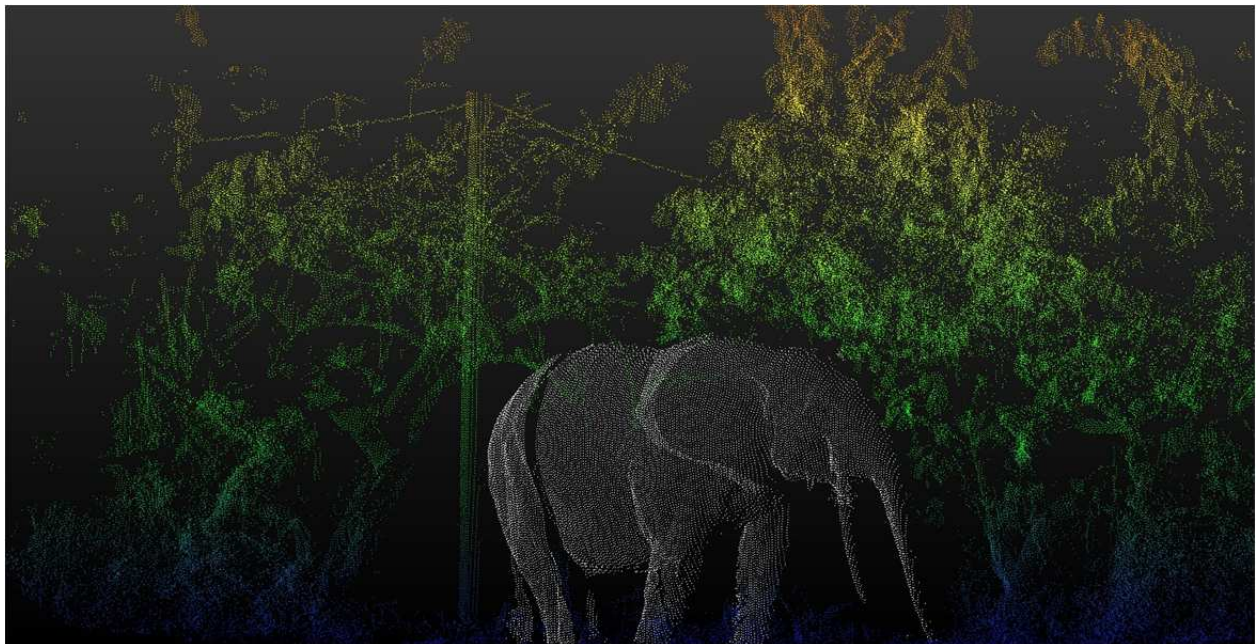
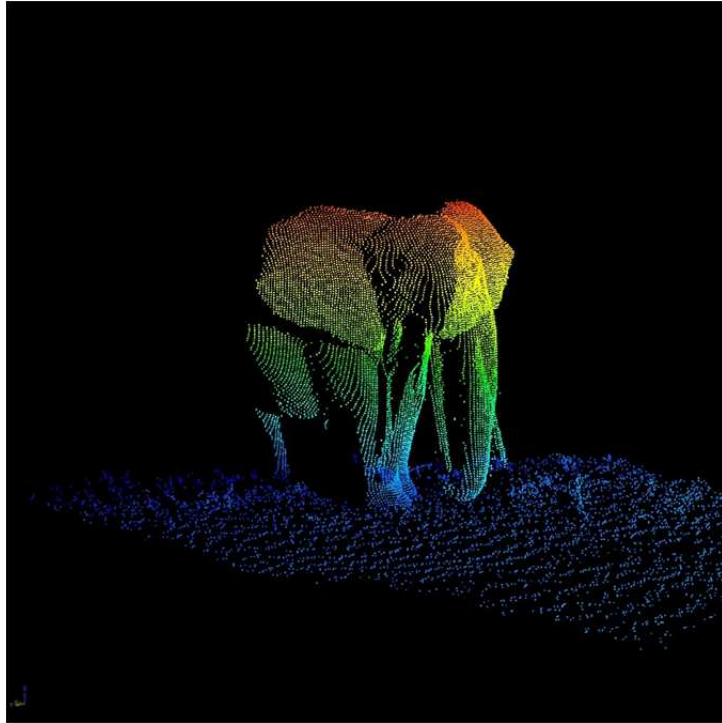
Predicted composition of all sites consistent with patterns of α and β



Fuzzy classification of grassland vegetation in an alkaline grassland in Püspökladány, Hungary, based on airborne LIDAR. Colours represent the weighted probability for a given vegetation class in each cell (0.5m²) (photo credit: András Zlinszky).



Vegetation composition of a peatland using Partial Least Square Regression models on a hyperspectral image. The image is a false colour composite showing the predicted abundance of Graminoids (Red), Shrubs (Green), and Bryophytes (Blue) (photo credit: Beth Cole).



A forest elephant “scanned” during a terrestrial laser-based measurement of a tropical rainforest in Gabon 2013 (photo credit: Kim Calders).